

Amendments to the Claims

This listing of claims will replace all prior versions and listings of claims in the application:

Claims 1-11 (cancelled)

- 12. (Currently amended) A method for generating a secondary library of <u>secondary</u> protein variants of a target protein comprising:
- a) inputting the three dimensional coordinates of said target protein into a computer;
- b) determining a criteria for including favorably ranked primary variant proteins for a primary library;
- bc) utilizing a force_field calculation to generate a-said primary library comprising a plurality of favorably ranked primary variant proteins comprising primary-variant amino acid residues at primary variant positions;
- d) determining a criteria for selecting amino acid residues from each of said variant positions from said favorably ranked primary variant proteins
- e) selecting amino acid residues from a plurality of said variant positions from said favorably ranked primary variant proteins; and
- ef) combining a plurality of said primary variant selected amino acid residues from step-b) to generate a said secondary library of said secondary variant proteins, wherein at least one of said secondary variant proteins is different from the said primary variant proteins.
- 13. (Previously presented) A method according to claim 12, wherein said force field calculation is a Self-Consistent Mean Field (SCMF) calculation.

Claims 14-20 (Cancelled)

- 21. (Currently amended) A method according to claim 12, further comprising synthesizing a plurality of said secondary variant proteins, wherein said combining comprises:
- eg) generating a set of oligonucleotide probes each encoding at least one of said primary variant amino acid residues at said variant positions;
- (h) using said probes in a polymerase chain reaction (PCR) to generate a plurality of oligonucleotide sequences, each encoding said secondary variant sequences; and (gi) producing said secondary variant sequences in host cells transformed with said
- oligonucleotide sequences.

- 22. (Previously presented) A method according to claim 21 wherein said PCR is multiple PCR wherein said probes are pooled.
- 23. (Previously presented) A method according to 22 wherein said probes are added in equimolar amounts.
- 24. (Currently amended) A method according to claim 22 wherein said probes are combined in amounts that correspond to the frequency of the said variant amino acid residues at said variant positions in said secondary library.

Claims 25-32 (cancelled)

- 33. (Currently amended) A method for generating a secondary library of <u>secondary</u> protein variants of a target protein comprising:
- a) inputting the three dimensional coordinates of said target protein into a computer;
- b) determining a criteria for including favorably ranked primary variant proteins for a primary library;
- bc) utilizing a force_field calculation to generate a primary library of favorably ranked primary variant proteins comprising a plurality of primary variant amino acid residues at primary variant positions;
- d) determining a criteria for selecting amino acid residues from each of said variant positions from said favorably ranked primary variant proteins
- e) selecting amino acid residues from a plurality of said variant positions from said favorably ranked primary variant proteins; and
- ef) combining a plurality of said primary variant selected amino acid residues from a plurality of said variant positions from said favorably ranked primary variant proteins step b) to generate a secondary library of secondary variant proteins.
- 34. (Currently amended) A method for generating a secondary library of <u>secondary</u> protein variants of a target protein comprising:
- a) inputting the three dimensional coordinates of said target protein into a computer;
 b) determining a criteria for including favorably ranked primary variant proteins for a primary library;